PATENT SPECIFICATION

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(54) NEW ESTERS OF SUBSTITUTED NICOTINIC ACIDS

We, THE BOOTS COMPANY LIMITED (formerly known as Boots Pure Drug Company Limited), a British Company, of 1 Thane Road West, Nottingham, England, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement: -

10 This invention relates to novel derivatives of nicotinic acid which have been found to

possess biological activity.

According to one aspect of the invention there are provided compounds of general 15 formulae I—IV

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II

III

IV

[Price 25p]

and pharmaceutically acceptable acid addition salts thereof in which

R₅ is phenyl;

(a) A_1 is ethylene and R_1 is dimethylamino, 1-pyrrolidinyl, piperidino or 4-

methyl-1-piperazinyl, or (b) A_1 is propylene and R_1 is dimethylamino, diethylamino, 1-pyrrolidinyl,

piperidino or morpholino;

A₂ is ethylene and R₂ is diethylamino, 1-pyrrolidinyl, morpholino or 4-methyl-1-piperazinyl, or

(d) A₂ is propylene and R₂ is dimethylamino, morpholino or 4-methyl-1piperazinyl;

A₃ is ethylene and R₃ is dimethylamino,

piperidino or morpholino, or

 A_3 is propylene and R_3 is dimethyl-

amino or 4-methyl-1-piperazinyl; and A_4 is ethylene and R_4 is dimethyl-(g) amino, or

 A_4 is propylene and R_4 is diethylamino,

1-pyrrolidinyl, piperidino or morpho-

The invention includes pharmaceutically acceptable acid addition salts of the compounds of general formulae I-IV. Typical salts falling within the invention include, for example, hydrochlorides, maleates, succinates and citrates. Details of many specific salts will be found in the examples at the end of this specification, but the acids used therein and which are listed above are only typical acids and are not intended to imply that the invention is limited to salts with these particular

Typical methods for the preparation of the compounds of the invention are as follows:-

[For the sake of brevity, the substituted pyridine nuclei of general formulae I-IV (less the —COO—A—R moeity) will be designated "B" from now on where convenient.]

(1) Trans-esterification of a compound of general formula V

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B-COOR

in which R_6 is C_{1-4} alkyl, preferably methyl or ethyl, with the required amino-alcohol of general formula VI

in which R represents R₁, R₂, R₃ or R₄ and A represents A₁, A₂, A₃ or A₄. This is carried out by heating such that the alcohol R₆OH which forms is readily eliminated by distillation as it is evolved during the reaction: -

R—A—OH . . . VI

$B-COOR_a+R-A-OH \Rightarrow B-COO-A-R+R_aOH$

In this way, and additionally by using an excess of the amino-alcohol as reaction medium, the equilibrium can be displaced towards the required product. Preferably a catalytic amount of sodium should be present.

The temperature required to achieve the desired result and the length of time of heating will naturally vary to some extent with the different values of B-COOR, and R—A—OH, but, in general, a temperature of at least 70° C. for at least 2 hours is advisable. For preference, to speed up the reaction and to ensure maximum yields, temperatures of the order of 120-180° C. are used for periods of 5—9 hours.

(2) Reaction of an acid chloride of general formula VII

B-COC1 . . . VII

with the required amino-alcohol of general formula VI hereinbefore described, optionally in an inert organic solvent such as benzene.

(3) Continuous azeotropic distillation with a neutral solvent boiling above 130° C. (preferably 130—160° C.) of a mixture of an acid of general formula VIII

B-COOH . . . VIII

and an amino-alcohol of general formula VI hereinbefore described. Examples of suitable solvents are xylene, chlorobenzene, ethylbenzene and cumene.

(4) Pharmaceutically acceptable salts of the bases prepared as described in (1)—(3) above are prepared by conventional methods. Thus, for example, a base may be dissolved in a suitable inert solvent such as a C1-4 alkenol (e.g. isopropanol) or tetrahydrofuran and the required acid added. Frequently the desired salt precipitates immediately or upon evaporation of some of the solvent; in other cases the addition of ether is necessary to cause precipitation of the salt.

The starting materials of the aforementioned general formulae V, VII and VIII are prepared by methods known in the art of pyridine chemistry.

It has been found that the compounds of the invention possess vasomotor properties viz. they are peripheral vasodilators, and may be used in the treatment of disorders of circulatory origin.

According to a further feature of the in-

vention there are provided therapeutic compositions which comprise a compound of the invention in association with pharmaceutical excipients for oral, rectal or parenteral administration. The compositions preferably contain 0.1-90% by weight of a compound of the invention.

Compositions for oral administration are the known pharmaceutical forms for such administration, such as for example tablets, capsules, syrups, and aqueous oily suspensions. The excipients used are the excipients known in the pharmacist's art. Thus, for example, tablets comprise a compound of the invention mixed with a conventional diluent such as lactose and a disintegrating agent such as maize starch and a lubricating agent such as magnesium stearate. Such tablets may if desired be provided with enteric coatings by known methods, for example by the use of cellulose acetate phthalate. Similarly capsules, for example hard or soft gelatin capsules, containing a compound of the invention, with or without other excipients, may be prepared by conventional means and, if desired, provided with enteric coatings. The tablets and capsules may conveniently contain 10-500 mg. of a compound of the invention.

Compositions for rectal administration are the known pharmaceutical forms for such administration, such as for example suppositories with cocoa butter or polyethylene glycol bases.

Compositions for parenteral administration, e.g. intravenous injection, are the known pharmaceutical forms for such administration, for example sterile solutions in normal saline for injection or sterile solutions in propylene glycol.

It will be appreciated that because of their physical characteristics (crystalline powders), the pharmaceutically acceptable acid addition salts hereinbefore described are to be preferred in most cases to the bases themselves (high boiling liquids).

The compositions hereinbefore described may be provided in dosage unit forms containing 70 mg.-14 g., more usually 140 mg.-1.4 g., optionally in divided dosage unit form.

Disorders of circulatory origin may be treated by a method comprising administering to a subject suffering from such disorders a peripheral vasodilating amount of a compound of the invention. Doses vary according to the activity of the particular compound, but in

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general fall within the broad range of 1-200 mg./kg., more usually within the range 2-20 mg./kg.

The following non-limitative examples

illustrate the invention.

Example 1.

3-N,N-Dimethylaminoropan-1-ol (13.4 g.) and sodium (0.07 g.) were added to a 100 ml. flask fitted with an inlet for dry nitrogen and provided with distillation means. After heating at about 50° C. until the sodium had dissolved, methyl 5-phenyl-6-methylnicotinate (10 g.) was added and heating continued for 9 hours at about 180° C.; methanol distilled off. After cooling, sodium amino-alcoholate was precipitated by the addition of dry ether (200 ml.) and filtered off. Evaporation of the ether and distillation of the residue in vacuo gave 3-N,N - dimethylaminopropyl 5 - phenyl-6-methylnicotinate, b.p. 145—150° C./0.01 mm.
The dihydrochloride was made by conventional means, m.p. 148° C. (isopropanol/ether).

By a similar technique, the compounds listed below were prepared.

m.p. Salt (°C) b.p. ester (°C./mm.) $\mathbf{A_2}$ Salt Et₂N---(CH₂)₂---165-170/0.01 dihydrochloride 146 190-195/0.02 180 178 200/0.1210-215/0.1 trihydrochloride 165 -(CH₂)₃---160 195/0.02 ,, 204 210-215/0.1 trihydrochloride ,,

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Example 2.

The apparatus and procedure of Example 1 were used, employing 3-N,N-dimethylamino-propan-1-ol (15.5 g.), sodium (0.07 g.), ethyl 2,6-dimethyl-5-phenylnicotinate (12.75 g.), and a temperature of 170—180° C. for about 8

hours. There was thus obtained 3-N,N-dimethylaminopropyl 2,6 - dimethyl - 5 - phenylnicotinate, b.p. 160—170° C./0.15 mm.; citrate, m.p. 70° C. (isopropanol/ether).

The compounds listed below were similarly

prepared.

CH ₃ CH ₃ COO-A ₃ -R ₃				
R_3	A_3	b.p. ester (°C./mm.)	Salt	m.p. salt (°C.)
Me ₂ N—	—(CH ₂) ₂ —	160—170/0.1	disuccinate	120
	. cc	150—160/0.05	disuccinate	136
○ -	22	190—195/0.15	dihydrochloride	145
Me-IV N-	—(CH ₂) ₃ —	210—215/0.1	trisuccinate	110

Example 3. The apparatus and procedure of Example 1 were used, employing 2-N,N-dimethylamino-- 15 ethan-1-ol (9 ml.), sodium (0.07 g.), methyl 5-phenylnicotinate (4.1 g.), and a temperature of 120—125° C. for 7.5 hours. The crude 2-N,N-dimethylaminoethyl 5-phenylnicotinate obtained as an oil was not distilled, but was used directly for the preparation of the maleate, m.p. 126° C. (tetrahydrofuran/ether).

The compounds listed below were similarly prepared.

T ₅ C00-A ₁ -R ₁			
R ₁	A_1	Salt	m.p. salt (°C)
	—(CH ₂) ₂ —	maleate	115
<u></u>	20	5 9	120
Me-N N-	33	trihydrochloride	200
Me ₂ N—	—(CH ₂) ₃ —	maleate	99
Et ₂ N—	93	22	144
	33		104
<u></u>	22	33	91
◇ -)	22	69

Example 4. to that of Example 1 the compounds listed below are prepared.

соо-А _Д -Ед
M CH ³

R ₄	A_4	b.p. ester (°C./mm.)	Salt	m.p. salt (°C.)
Me ₂ N—	—(CH ₂) ₂ —	65/0.09	maleate	91
Et ₂ N—	—(CH ₂) ₃ —	65/0.09 116—120/0.8	>>	99
	>>	140—142/0.3	22	96
<u></u> -))	128/0.05	2)	118
◆-	22	130/0.05	2)	120

[All the compounds of the invention described in Examples 1—4 gave satisfactory elemantal analyses and their structures have been verified by infra-red spectroscopy.]

Example 5.

In the preparation of tablets, mixtures of the following type may be tableted in conventional manner:

10	Pharmaceutically acceptable	
	salt of the invention	10—90%
	Lactose	080%
	Maize starch	5—10%
	Magnesium stearate	ca.1%
15	Microcrystalline cellulose	0—90%
	•	(by weight)

Example 6.

In the preparation of capsules, a salt of the invention may be mixed with an equal weight of lactose and the mixture encapsulated in hard gelatin capsules.

Example 7. In the preparation of 1 g. suppositories, bases of the following type may be used, each suppository containing for example 200 mg. of salt of the invention: -

Polyethylene glycol Polyethylene glycol	33% 47%
Water	20%

Example 8.

Solutions for parenteral injection may be prepared comprising 4 mg. of a salt of the invention per ml. of normal saline for injection B.P.

WHAT WE CLAIM IS: -1. Compounds of general formulae I—IV

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II

 \mathbf{III}

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7 1,352,415 7 12. 2 - Piperadinoethyl 2,6 - dimethyl - 5and pharmaceutically acceptable acid addition phenylnicotinate and pharmaceutically acceptsalts thereof in which able acid addition salts thereof. R_5 is phenyl; 13. 2 - Piperidinoethyl 2,6 - dimethyl - 5-(a) \bar{A}_1 is ethylene and R_1 is dimethylamino, 65 5 phenylnicotinate disuccinate. 1-pyrrolidinyl, piperidino or 4-methyl-14. 2 - Morpholinoethyl 2,6 - dimethyl - 5-1-piperazinyl, or phenylnicotinate and pharmaceutically accept-A₁ is propylene and R₁ is dimethylable acid addition salts thereof. amino, diethylamino, 1-pyrrolidinyl, 15. 2 - Morpholinoethyl 2,6 - dimethyl - 5piperidino or morpholino; 70 A₂ is ethylene and R₂ is diethylamino, phenylnicotinate dihydrochloride. 10 16. Compounds as claimed in claim 1 and 1-pyrrolidinyl, morpholino or 4-methylof general formula IV in which (a) A4 is 1-piperazinyl, or ethylene and R₄ is dimethylamino, or (b) A₄ A₂ is propylene and R₂ is dimethylis propylene and R₄ is diethylamino, 1-pyrroliamino, morpholino or 4-methyl-1-75 dinyl or piperidino. 15 piperazinyl; 17. 3-Piperidinopropyl 2-methylnicotinate A₃ is ethylene and R₃ is dimethyland pharmaceutically acceptable acid addition amino, piperidino or morpholino, or A₃ is propylene and R₃ is dimethylamino or 4-methyl-1-piperazinyl; and salts thereof. 18. 3-Piperidinopropyl 2-methylnicotinate 80 A₄ is ethylene and R₄ is dimethylmaleate. 20 19. 3-Piperidinopropyl 2-methylnicotinate. amino, or A_4 is propylene and R_4 is diethylamino, 20. 3 - (Pyrrolidin - 1 - yl)propyl 2 - methyl-(h) nicotinate and pharmaceutically acceptable 1-pyrrolidinyl, piperidino or morphoacid addition salts thereof. 21. 3 - (Pyrrolidin - 1 - yl)propyl 2 - methyl-25 2. Compounds as claimed in claim 1 and of general formula I in which (a) A1 is ethylene nicotinate maleate. 22. A process for preparing the compounds and R₁ is dimethylamino, 1-pyrrolidinyl or claimed in any one of claims 1-21 substanpiperidino, or (b) A₁ is propylene and R₁ is dimethylamino, diethylamino, 1-pyrrolidinyl, tially as described herein. 23. Therapeutic compositions which compiperidino or morpholino. prise as an active ingredient a compound as 3. 2-(Pyrrolidin-1-yl)ethyl 5-phenylnicotinclaimed in any one of claims 1-21 in associaate and pharmaceutically acceptable acid addition with a pharmaceutical excipient for oral, tion salts thereof. rectal or parenteral administration. 4. 2-(Pyrrolidin-1-yl)ethyl 5-phenylnicotin-24. Compositions as claimed in claim 23 in 35 ate maleate. 5. 3 - Piperidinopropyl 5 - phenylnicotinate the form of tablets or capsules. 25. Compositions as claimed in claim 23 in and pharmaceutically acceptable acid addition the form of syrups, aqueous suspensions or oily salts thereof. 6. 3-(Pyrrolidin-1-yl)propyl 5-phenylnicotinsuspensions. 26. Compositions as claimed in claim 23 in 100 ate and pharmaceutically acceptable acid addi-

tion salts thereof. 7. Compounds as claimed in claim 1 and of general formula II in which (a) A2 is ethylene and R2 is 1-pyrrolidinyl, morpholino 45 or 4-methyl-1-piperazinyl, or (b) A2 is propylene and R₂ is dimethylamino, morpholino

or 4-methyl-1-piperazinyl. 8. 2 - (4 - Methylpiperazin - 1 - yl)ethyl 6methyl-5-phenylnicotinate and pharmaceuti-50 cally acceptable acid addition salts thereof.

9. 2 - (4 - Methylpiperazin - 1 - yl)ethyl 6methyl-5-phenylnicotinate trihydrochloride.

10. 2 - (Pyrrolidin - 1 - yl)ethyl 6 - methyl-5 - phenylnicotinate and pharmaceutically 55 acceptable acid addition salts thereof.

11. Compounds as claimed in claim 1 and of general formula III in which (a) A3 is ethylene and R₃ is piperidino or morpholino, or (b) A3 is propylene and R3 is dimethylamino or 4-methyl-1-piperazinyl.

the form of suppositories. 27. Compositions as claimed in claim 23 for parenteral administration.

28 Compositions as claimed in claim 27 in the form of solutions.

29. Compositions as claimed in any one of claims 23-28 in which the active ingredient is in the form of a pharmaceutically acceptable acid addition salt.

30. Compositions as claimed in any one of claims 23-28 in which the active ingredient is 3-piperidinopropyl 2-methylnicotinate or a pharmaceutically acceptable acid addition salt

31. Compositions as claimed in any one of claims 23-28 in which the active ingredient is 2-(pyrrolidin-1-yl)ethyl 5-phenylnicontinate or a pharmaceutically acceptable acid addition salt thereof.

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